Vascular Resistance Characteristics of 7,12-Dimethylbenz(a)anthracene-induced Rat Mammary Tumors and Normal Tissues as Studied in Vitro

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ABSTRACT

Vascular perfusion characteristics have been studied in dimethylbenz(a)anthracene-induced rat mammary neoplasia and compared with those of skin, skeletal muscle, salivary gland, kidney, spleen, uterus, and brain by means of an artificial perfusion technique. Perfusion of tissues and organs was measured by the microsphere tracer technique. This procedure makes possible a detailed hemodynamic analysis of several tissues under controlled conditions, in this study maximum vascular relaxation, without confounding endogenous vasoregulation. The maximal perfusion capacity, i.e., during smooth muscle relaxation, of tumors and various tissues was related to perfusion pressure at three levels by means of three differently labeled microspheres. Tumors, especially large ones, have a low maximum perfusion capacity, i.e., high vascular resistance, compared to most other tissues. For the tumors, a relatively high perfusion pressure is required to open up the otherwise collapsed vascular network.

INTRODUCTION

Contradictory results on total blood flow and vascular reactivity of tumor tissue as compared to various normal tissues can be ascribed partly to the use of different tumor models and methods of analysis (cf. Ref. 1). Most studies are, however, performed in vivo, either during anesthesia or under conscious conditions. The central and peripheral hemodynamics thus vary greatly (14), complicating the interpretation of data obtained. The vascular tone in various organs and tissues also varies greatly within the same animal at the time of experimentation, and no base-line vascular tone to which the tumor vasculature can be related can be achieved.

An in vitro perfusion technique was therefore developed as a further elaboration of the procedure described by Folkow et al. (2). In light of recently published data on blood flow in 7,12-dimethylbenz(a)anthracene-induced mammary tumor of the rat (5) and findings of an increased interstitial fluid pressure within these tumors (11), it was considered of interest to elucidate the maximal perfusion capacities in these tumors in relation to various perfusion pressures and to compare these results with findings in various normal tissues.

MATERIALS AND METHODS

Tumor Model. Female Sprague-Dawley rats (Anticimex, Stockholm, Sweden), 50 to 55 days old, were fed by gavage with 7,12-dimethylbenz(a)anthracene, 16 mg dissolved in 1 ml of olive oil (4, 13). Multiple mammary tumors became overt from the sixth week after induction, and experiments were performed 4 weeks later.

Perfusion Technique. The rats were anesthetized with Nembutal (50 mg/kg body weight i.p.). The abdomen was opened in the midline by cautORIZATION, the stomach and intestines were removed, and the vessels and cut ends of viscera were carefully ligated. The thorax was cut open by cautORIZATION, and the aortic root was rapidly connected to a pump system. The right cardiac ventricle was cut open, a cannula was inserted for free outflow, and the perfusion was started. The caudal and one femoral artery were cannulated for measurement of pressure and reference flow, respectively. The femoral vein was cannulated for measurement of venous pressure. Pressures were continuously recorded on a Grass polygraph. The oxygenated perfusate, kept at 38°C, consisted of 6% dextran (Macrodex, AB Pharmacia, Upssala, Sweden; mean M, 70,000) and 100 ml of horse serum (normal serum; SBL, Stockholm, Sweden) in 1,000 ml of a salt solution [Na+, 143 mM; K+, 4.3 mM; Ca2+, 2.5 mM; Mg2+ 0.83 mM; Cl-, 141 mM; HCO3; 13.3 mM; H2PO4, 0.46 mM; and glucose, 5.6 mM (9)]. The perfusion system is illustrated in Chart 1. The pump used was a peristaltic constant-flow type Ismatec MP 4. To monitor possible fluid retention, the animal was placed on a balance throughout the experiment. The temperature of the preparation was kept constant at 37°C by a heating lamp mastered from a rectal thermometer. Papaverine was used to obtain maximal smooth muscle relaxation. When no further relaxation could be obtained by repeated papaverine injection, the flow was set to various levels, and arterial and venous pressures were measured. In this way, a flow-pressure relationship was obtained for the whole preparation.

Regional Blood Flow Determination. At 3 different flow-pressure levels, regional blood flow was determined by injection of polystyrene spheres (3M Co., St. Paul, MN), with diameters of 15 ± 3 μm (SD), into the tubing connected to the aortic root. The spheres were labeled with 141Ce, 85Sr, and 51Cr, respectively, and were given approximately 150,000 at a time. During and immediately after microsphere injection (90 s), a reference perfusate sample was drawn from the femoral artery at a rate of 0.3 ml/min.

After the perfusion experiment, the tumors and parts of the quadriceps muscle, the paw, spleen, brain, salivary gland, uterus, and kidneys were dissected out, weighed, and placed in vials for activity measurement in a Packard Auto-Gamma spectrometer Model 5019. From the activity in the reference samples, perfusate flow in the tissue samples could be calculated (6).

Care was taken to perform the experiments in a standardized way and as rapidly as possible, since prolongation leads to edema and increased venous pressure. Experiments exhibiting such artifacts were rejected.

Statistics. Analyses were performed according to Student's t test using a pairing design. Significant differences from the tumor pressure-flow characteristics are indicated by * (P < 0.05), ** (P < 0.01) and *** (P < 0.001) in Chart 1.

RESULTS

Six successful perfusion experiments were performed, but a larger number of experiments were rejected due to technical failures resulting in edema, increased venous pressure, or uneven distribution of microspheres between the 2 kidneys. In the 6 animals, 24 tumors were analyzed, with a mean weight of 2.9 g. The perfusate flow to the animals was set to produce an initial...
perfusion pressure (arterial pressure minus venous pressure) of 13.4 ± 0.5 mm Hg, an intermediate pressure of 24.9 ± 0.7 mm Hg, and a final pressure of 34.2 ± 1.6 mm Hg. Absolute perfusate flow data were obtained for each of the above-mentioned tissues in each animal for each of the 3 pressure levels. Pressure-flow data are summarized in Chart 2, from which it will be seen that the flow capacity of tumors is low compared to most other tissues.

Organ vascular resistance is calculated by dividing the perfusion pressure by the perfusate flow, and this measure is considered to represent the functional state of the vascular beds in organs and tissues more adequately. It will be seen from Chart 3 that the vascular resistance is high in tumors and that the resistance seems to decrease more rapidly with increased perfusion pressure in tumors than in most other organs. Another way to express this concept is to present the relative increase in assumed cross-sectional area of the vascular beds, this area being inverse to the square root of the vascular resistance according to Poiseuille's law. These data, related to distending pressure (the mean of the arterial and venous pressure), are presented in Chart 4, from which it will be seen that the cross-sectional area of the tumor vascular bed increases more rapidly than in other tissues.

DISCUSSION

This study was undertaken to elucidate 2 aspects of tumor vascular functional morphology and physiology not possible to study in the living animals, i.e., the maximal perfusion capacity during standardized maximal vasodilation of all vascular beds representing the available capillary cross-sectional area and the characteristics of the pressure-flow curves at low perfusion pressures.

In vivo, the various regional vascular beds are under undefined and different vasoconstrictor tone, even under so-called resting conditions in the conscious or anesthetized animal. This makes reliable estimation of the blood flow capacity in various tissues under identical vascular constriction, and thus the relative available cross-sectional area of the vascular beds in various tissues, impossible.

An in vitro perfusion model was therefore considered useful for the present purposes. A similar perfusion system was used to characterize the vascular bed of rat hind quarters (2). However, only bulk flow as derived from the pump settings was...
that maximal vasodilation is rarely present under in vivo conditions, thus exposing the capillary bed to this pressure in full. The maximal perfusion capacity at any perfusion pressure is low for tumor tissue. In vivo, measurement of resting blood flow in this animal model (5) showed comparatively high flow values in tumor tissue compared to various other tissues. This indicates that the tumor vascular bed is probably maximally dilated under resting conditions, in contrast to most other vascular beds. The unstrained exposure of the tumor capillary bed to a high blood perfusion pressure might result in edema and increased interstitial fluid pressure, which was actually recorded in these tumors (11). Recently published findings of decreasing interstitial fluid pressure within these tumors upon noradrenaline infusion further substantiate this concept (10).

The curves in Chart 4, showing the relative cross-sectional area of the vascular beds in relation to distending pressure show that this area increases more rapidly for a certain increment in distending pressure in tumor tissue than in the other vascular beds studied. This finding can probably be attributed to an extravascular, interstitial tissue pressure counteracting the intra-vascular distending pressure. Thus, the "critical closing pressure" (7), not studied here due to technical difficulties in measuring perfusate flow at perfusion pressures below 13 mm Hg, is probably higher in tumor tissue than elsewhere. This finding indicates an impact on vascular perfusion of the increased interstitial fluid pressure recently recorded in these tumors (11) as well as in other tumors (3, 8). The hypothesis of a "compartment syndrome" taking part in the development of tumor hypoxia and necrosis is substantiated.

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